

sion, stated that “the standard of care identified in this article has not been imposed by a court of law. It is the standard of care developed by geneticists and physicians. Debate as to its ‘reasonableness’ will have to be resolved by the medical genetics community” (Sharpe 1994a).

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References

- Ad Hoc Committee on Genetic Counseling (1975) Genetic counseling. *Am J Hum Genet* 27:240–242
- Almqvist E, Adam S, Bloch M, Fuller A, Welch P, Eisenberg D, Whelan D, et al (1997) Risk reversals in predictive genetic testing for Huntington disease. *Am J Hum Genet* 61:945–952
- American Board of Medical Genetics Inc (1998) Bulletin of information—description of examinations. National Board of Medical Examiners, Bethesda, MD
- Canadian College of Medical Geneticists (1998) Training guidelines for genetic centres and candidates for fellowship. Canadian College of Medical Geneticists, Ottawa
- De Wert G (1992) Predictive testing for Huntington disease and the right not to know: some ethical reflections. *Birth Defects* 28:133–138
- Fitzpatrick JL, Hahn C, Costa T, Huggins MJ (1999) The duty to recontact: attitudes of genetics service providers. *Am J Hum Genet* 64:852–860
- Jadad AR, Gagliardi A (1998) Rating health information on the Internet: navigating to knowledge or to Babel? *JAMA* 279:611–614
- Kim P, Eng TR, Deering MJ, Maxfield A (1999) Information in practice: published criteria for evaluating health related web sites. *BMJ* 318:647–649
- National Society of Genetic Counselors (1997) Predisposition genetic testing for late-onset disorders in adults. *JAMA* 278:1217–1220
- Ost D (1984) The “right” not to know. *J Med Philos* 9:301–312
- Sharpe NF (1994a) Informed consent and Huntington disease: a model for communication. *Am J Med Genet* 50:239–246
- (1994b) Psychological aspects of genetic counseling: a legal perspective. *Am J Med Genet* 50:234–238
- Tresemmer v Barke (1978) 86 Cal App 3d 656; 150 Cal Rptr 384
- Yarborough M, Scott JA, Dixon LK (1989) The role of beneficence in clinical genetics: non-directional counseling reconsidered. *Theor Med* 10:139–149

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Reply to Sharpe

To the Editor:

Mr. Sharpe correctly notes that in our article (Fitzpatrick et al. 1999) he was credited for considering the existence of a physician’s duty of care toward patients (Sharpe 1994). His comments in this regard were indeed made in the context of Huntington disease, but, as we did not attribute to him *any* opinion on the duty to recontact, his position on this subject was not misrepresented, but simply omitted, from our discussion. We apologize to Mr. Sharpe and thank him for clarifying his position. The intention of our article was to report and discuss original research findings and not to present a detailed analysis of medical principles and legal obligations associated with a theoretical duty to recontact. It was our hope that our article would stimulate such a discourse, and we thank Mr. Sharpe for his insightful comments.

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References

- Fitzpatrick JL, Hahn C, Costa T, Huggins MJ (1999) The duty to recontact: attitudes of genetics service providers. *Am J Hum Genet* 64:852–860
- Sharpe NF (1994) Psychological aspects of genetic counseling: a legal perspective. *Am J Med Genet* 50:234–238

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The Choice to Have a Disabled Child

To the Editor:

What are the purposes of genetic testing, what are the principles guiding its use, and who should decide what tests should be available for what purposes? These familiar questions are raised in an unfamiliar context by a study reported recently in this journal (Middleton et al. 1998). Attitudes toward genetic testing were assessed among deaf people attending a conference in the United